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PATENT ABSTRACTS OF JAPAN, vol. 9, no. 175 (C-292)[1898], 19th July 1985; & JP-A-60 47 053

PATENT ABSTRACTS OF JAPAN, vol. 14, no. 342 (C-743)[4285], 24th July 1990; & JP-A-2 124 955

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Description

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BACKGROUND OF THE INVENTION

This invention relates to an extrudable adhesive composition of ethylene copolymer and tacklifying resin, useful for bonding structural polymers to barrier polymers.

Many attempts have been made to provide adhesive compositions which are particularly suitable for joining together layers of polymers, especially for joining structural polymers to polymers such as barrier resins. For example, Japanese Application 53 18653 discloses an adhesive resin for low temperature heat sealing purposes comprising 3 to 100 parts by weight of a hydrogenated petroleum resin, and 100 parts by weight of an ethylene carboxylate-vinyl acetate copolymer obtained by copolymerizing an unsaturated carboxylic acid and/or anhydride with at least one vinylidene monomer in the presence of an ethylene-vinyl acetate copolymer.

Japanese application 60 51768 discloses a hot melt adhesive composition of 100 parts by weight of polymer component, 25 to 200 parts by weight of tackifier resin, and 75 parts by weight or less of wax. The polymer component comprises 25 to 75 weight % low crystallinity or amorphous ethylene alpha-olefin copolymer (20 to 50 mole % ethylene, 50 to 80 mole % alpha-olefin) and 25 to 75 weight % ethylene vinyl ester copolymer. Either polymer, or both, are modified with unsaturated carboxylic acid or its acid anhydride. The tackifier resin is a terpene resin or petroleum resin. The wax is a paraffin, microcrystalline polyethylene, etc. The composition has a viscosity of 50 Pa.sec or less at 180 °C.

Japanese Application 01 27923 discloses a laminate in which a resin layer is coextrusion laminated onto an oriented film or Al foil. The adhesive is made of 50 to 99 weight percent ethylene-alpha-olefin random copolymer partially graft-modified with acid, and 1 to 50 weight percent tackifier. The ethylene-alpha-olefin copolymer has a density of 0.850 to 0.900 g/cc.

U.S. Patent 4,670,349 discloses an adhesive resin composition comprising 100 parts of an ethylene/vinyl acetate or ethylene/alpha-olefin random copolymer, 1-50 parts of modified polyethylene containing 0.01 to 10% grafted acid or derivative, and 1-125 parts hydrogenated aromatic petroleum resin. The resin composition is useful as an adhesive for polystyrene and ethylene vinyl alcohol copolymer.

U.S. Patent 4,337,297 discloses blends of a low density ethylene polymer and a copolymer of 70-90 % ethylene with the balance acrylic or methacrylic acid or vinyl acetate, the two ethylene polymers comprising together 100 parts; an ionomer resin having an acid content of 1-10%, 2-100 parts; and optionally a tackifying resin, 0-150 parts. The blend is a useful adhesive for PET film.

U.S. Patent 4,367,113, Karim et al. discloses a similar blend except the low density polymer is replaced by an ethylene-alpha olefin copolymer or isotactic propylene polymer.

U.S. Patent 4,358,557 discloses a hot-melt adhesive comprising ethylene vinyl acetate copolymer, a tackifier resin selected from hydrocarbon resin, polyterpene resin or rosin ester, a paraffin wax and an aromatic hydrocarbon resin.

U.S. Patent 4,861,676 discloses a blend of ethylene copolymer, ethylene copolymer modified by grafting with a comonomer containing pendant acid or acid derivative functionality, styrene polymer, and alicyclic resin modifier or stabilized rosin ester. The composition provides a useful adhesive, particularly for bonding polystyrene to barrier polymers.

U.S. Patent 4,861,677 discloses a blend of ethylene vinyl acetate copolymer, ethylene vinyl acetate copolymer modified by grafting with a comonomer containing pendant acid or acid derivative functionality, and impact-modified polystyrene. The composition provides a useful adhesive, particularly for bonding polystyrene to gas barrier polymers.

SUMMARY OF THE INVENTION

The present invention provides an extrudable bonding resin composition consisting essentially of:

(a) about 65 to about 99 percent by weight of an ethylene copolymer portion consisting essentially of (i) about 1 to 100 percent by weight of a copolymer of about 50 to about 80 weight percent ethylene and 20 to about 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and optionally carbon monoxide, the carbon monoxide being present in an amount of 0 to about 30 weight percent; having grafted thereon side chains at least one comonomer unit in length, the comonomer units in said side chains being selected from the group consisting of ethylenically unsaturated mono-, di-, or polycarboxylic acids, ethylenically unsaturated

carboxylic acid anhydrides, metal salts and half-esters of such acids and anhydrides, and mixtures thereof, wherein the amount of said grafted comonomer units comprises about 0.03 to about 10 percent by weight of the total bonding resin composition, and

(ii) 0 to about 99 percent by weight of at least one copolymer of about 50 to about 80 weight percent ethylene and 20 to about 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and optionally carbon monoxide, the carbon monoxide being present in an amount of 0 to about 30 weight percent,

wherein the copolymer of (i) and each copolymer of (ii) contain at least one said copolymerized comonomer in common, the amount of each such common comonomer in each copolymer of (ii) being within about 10% of the amount of the corresponding comonomer in copolymer (i), and the total amount of said copolymerized comonomers other than such common comonomers in any such copolymer being less than about 10%; and

(b) about 1 to about 35 percent by weight of a tackifying resin.

The invention further provides a multiple layer structure comprising at least one structural layer, at least one barrier layer, and at least one bonding layer of the above extrudable bonding resin composition.

DETAILED DESCRIPTION OF THE INVENTION

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The extrudable bonding resin of the present invention comprises a blend of about 65 to about 99 percent by weight, preferably about 80 to about 95 percent by weight, of an ethylene copolymer component, and about 1 to about 35 weight percent of a tackifying resin. The ethylene copolymer component comprises an ethylene copolymer grafted with pendant acid or certain acid derivative functionality. This grafted copolymer may be blended, if desired, with additional ungrafted ethylene copolymer. Such blending may be desirable in order to minimize the amount of the relatively more expensive grafted material, while maintaining the excellent adhesive properties of the composition.

The grafted ethylene copolymer is present in amounts of about 1 to about 100, and preferably about 5 to about 50 percent by weight of the ethylene copolymer portion of the composition. This component is a copolymer containing about 20 to about 50, and preferably about 25 to about 30 weight percent copolymerized comonomer selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and carbon monoxide, the carbon monoxide being present in an amount of 0 to about 30 weight percent. (The amount of carbon monoxide is limited only because it is believed that preparation of polymers containing more than about 30 weight percent carbon monoxide is not feasible due to problems of reactor fouling.) Vinyl acetate is a preferred comonomer, but methyl acrylate, n-butyl acrylate, and the like, optionally with carbon monoxide present as a third monomer, are also quite suitable. The balance of the copolymer is substantially copolymerized ethylene. Such copolymers are prepared by the well-known addition polymerization processes. Onto this copolymer are grafted additional comonomer units of carboxylic acid or certain carboxylic acid derivative functionality to form short sidechains. (It is believed that when maleic anhydride is the graft comonomer, the sidechains are only one or at most two monomer units long.) The melt index of the resulting graft copolymer, as measured by ASTM D1238 Condition "E", should be about 0.5 to about 40. Outside of these ranges, processing becomes more difficult, and flow instabilities may result.

The grafting monomer is selected from the group consisting of ethylenically unsaturated mono-, di-, or polycarboxylic acids, ethylenically unsaturated carboxylic acid anhydrides, and metal salts and half-esters of such acids and anhydrides. Examples of the acids and anhydrides include acrylic acid, methacrylic acid, maleic acid, fumaric acid, itaconic acid, crotonic acid, itaconic anhydride, maleic anhydride, and dimethyl maleic anhydride, and salts and half-esters thereof. Among the acids and anhydrides that are particularly useful are maleic anhydride and maleic acid.

The method for grafting of the comonomer onto the ethylene copolymer can be any of the processes which are well known in the art. For example, grafting can be carried out in the melt without a solvent, as disclosed in European Patent Application 0 266 994, incorporated herein by reference, or in solution or dispersion. Melt grafting can be done using a heated extruder, a Brabender™ or Banbury™ mixer or other internal mixers or kneading machines, roll mills, and the like. The grafting may be carried out in the presence of a radical initiator such as a suitable organic peroxide, organic perester, or organic hydroperoxide. The graft copolymers are recovered by any method which separates or utilizes the graft polymer that is formed. Thus the graft copolymer can be recovered in the form of precipitated fluff, pellets, powders, and

the like.

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The amount of monomer grafted onto the ethylene copolymer is not particularly limiting, and may be as low as about 0.03 weight percent or as much as about 5 percent or even 10 percent, based on the weight of the grafted ethylene copolymer. The total amount of graft polymer in the total adhesive composition, however, is important, and should be between about 0.03 and about 10 weight percent of the total composition, in order to obtain superior adhesive and bonding properties.

The ungrafted ethylene copolymer component comprises at least one comonomer similar to that described above, but without the grafted comonomer. In order to achieve good peel strength in laminates prepared from the present adhesive, the comonomer of the ungrafted copolymer should be of a type and amount sufficiently similar to that of the grafted copolymer that the grafted and ungrafted copolymers have a suitable degree of compatibility. Such copolymers are normally compatible if they contain the same or closely chemically related monomers (except for the graft comonomer) and the amounts of such monomers in the two copolymers are similar. Specifically, the grafted copolymer and each of the ungrafted copolymers (if there is more than one) should contain at least one said copolymerized comonomer in common and the amount of each such common comonomer in the grafted copolymer and the ungrafted copolymer should be within about 10%, and preferably within about 5%, of each other. Thus if the grafted copolymer is based on an ethylene polymer containing 30 weight % n-butyl acrylate, the ungrafted copolymer should contain between about 20 and about 40 weight percent n-butyl acrylate. Small amounts of additional comonomers may also be present in one or both copolymers, but in order to retain compatibility each copolymer should contain less than about 10% by weight of such comonomers that are not common to the other copolymer. Alternatively, closely similar comonomers, such as n-butyl acrylate and i-butyl acrylate could be interchanged while retaining compatibility. It is preferred that the graft copolymer be the same as the ungrafted copolymer, except for the presence of the grafted monomers. The melt index of the ungrafted copolymer should be about 0.05 to about 40.

The ungrafted copolymer component can be a single polymer or it can be more than one polymer, provided that the comonomer content of each such polymer is as described above, i.e., the polymers are mutually compatible. It has been sometimes found to be desirable that the ungrafted copolymer component be a mixture of two polymers having identical comonomer contents but having differing melt indices. For example, one copolymer can have a melt index of about 6 and a second can have a melt index of 25 or higher. The mixture of these polymers should have an overall melt index within the range of about 0.05 to about 40, as indicated. Use of such a mixture of copolymers provides ease in processability in, for example extrusion coating.

It is similarly to be understood that the use of more than one graft copolymer will fall within the scope of the present invention, provided that the general principles of polymeric compatibility set forth herein are maintained.

The second major component of the present invention is a tackifying resin, which is present at about 1 to about 35 weight percent, preferably about 5 to about 20 weight percent of the composition. Suitable tackifying resins may be selected from the categories of:

- (i) alicyclic or aliphatic hydrocarbon resins;
- (ii) aromatic hydrocarbon resins;
- (iii) rosin and rosin derivatives; and
- (iv) terpene resins, or mixtures thereof.

These tackifying resins will generally have a ring-and-ball softening temperature (ASTM E-2858T) of 0 to 150 °C, preferably about 75 to about 140 °C.

The tackifying resins employed in the present invention are well known products available from commercial sources. Rosin tackifiers are described in the Kirk Othmer Encyclopedia of Chemical, Technology, Interscience Publishers, Second Edition, Volume 17, pages 475-509. They include naturally occurring rosins and chemically modified rosin derivatives obtained by hydrogenation, dehydrogenation, isomerization, and the like. Rosin derivatives includes rosin esters and rosin acids. Rosin acids are typically derived from tall oil and can be mixtures of so called abietic types and primary types. Rosin esters are formed by esterifying rosin acid with a di-, tri-, or tetra-hydroxy alphatic alcohol such as ethylene glycol, propylene glycol, glycerine, or pentaerythritol. The terpene resins are generally prepared by the polymerization of terpene hydrocarbons in the presence of Friedel-Crafts catalysts at moderately low temperatures. Petroleum resins, under which are classed aliphatic, alicyclic, and aromatic hydrocarbon resins, are described in the Kirk Othmer Encyclopedia of Chemical Technology, Interscience Publishers, Third Edition, Volume 12, page 852. They are generally prepared by polymerization of 4-10 carbon atom hydrocarbons by selected Friedel Crafts catalysts. Higher or lower hydrocarbons may also be present. The product may be further partially or fully hydrogenated. Suitable aromatic resins can be prepared from polymerization of alpha methyl styrene,

vinyl toluene, and/or indene monomers.

In addition to the above mentioned components, the adhesive resin may contain small amounts of other materials commonly used and known in the art, such as antioxidants, stabilizers, slip additives, and fillers. The adhesive resin composition is prepared by blending the above described components by any suitable means, such as melt blending, extruding, etc. Further details are provided in the Examples which follow.

The composition of the present invention provides excellent adhesion to polyester and copolyester resins, vinyl chloride polymers, polycarbonates, polyethylenes, ethylene copolymers, ionomers, polystyrene, ethylene vinyl alcohol copolymer, polyamides, and the like. It also shows good adhesion to substrates such as aluminum foil. The adhesive is particularly useful for extrusion or coextrusion laminating structural and barrier polymers on to substrates such as polyester film, including multiple layer structures comprising at least two structural layers and at least one barrier layer. The structural layers may be of the same material or can be of different materials, e.g. one layer polyester and a second layer ionomer or acid copolymer. The adhesive can be used without having to resort to use of a solvent-based primer to prepare the polymer film surface. The resulting composite structures are useful in packaging and other applications.

Examples

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Adhesive blends for the Examples shown in Table 1 were prepared by dry blending the ingredients in a polyethylene bag and subsequently melt blending at 210 to 230 °C in a 28 or 53 mm Werner & Pfleiderer™ twin screw extruder with a vacuum port. Each of the blends, listed in the Table, also contained about 0.10 weight percent Irganox™ 1010 hindered polyphenol stabilizer (not separately reported in the Table). For the extruded pellets, 0.3 weight percent Kemamide™ W20 was also added.

The adhesive properties of the blends were evaluated by measuring the peel strength of laminates prepared using a sheet of the adhesive between two layers of PET film. The layers of PET film were 12 micrometers (0.48 mils) thick; heat seal conditions were 210 °C, 0.8 seconds dwell time and 324 kPa (47 psig) applied pressure. Peel strength was measured by ASTM D 1876-72, modified in that the test was run at 305 mm/min, rather than 254 mm/min, and 3 to 6 duplicates of each sample were measured, rather than 10. The numbers are given as grams per 25 mm.

Comparative Examples C1 through C8 show poor peel strength because the acid polymer, which contains 0% vinyl acetate comonomer, is used with a base resin containing 28% vinyl acetate. This is greater than the 10% difference in comonomer content between the ethylene polymer and its acid-modified ethylene polymer permitted in the present invention. In contrast, Examples 1 through 4 show good results because the difference in comonomer content between the acid polymer and the base resin is less than 10%.

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10		re & MAn		sters blend	0	0	•	0	0	0	0	0	0.1	0.30	0.7	0.30	ethylene with 28 weight percent vinyl	•	ethylene with 10	acrylate and 10 weight percent	a copolym	Ite, grait	100% hydrogenated C9 hydrocarbon resin,	having a Ring and Ball softening point	Pils is a similar material, R&B 115.C.
15		Adhesive	Thickness	micrometers	41	36	33	33	33	38	38	41	38	30	30	43	ith 28 We			nd 10 wei	/min, or	nyi aceto e. melt i	C9 hydro	d Ball sc	r materia
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30			Acid Copol.b	Type	E/10iBA/10MAA	ŧ	E	E	=	E	E	E	E/28VA-g-1.5MAn	£	2	r	A is a copolymer of	e, Be	Acid copolymer is either a terpolymer of	weight percent isobutyl	methacrylic acid, melt index 10 dg/min, or a copolymer	ethylene with 28 Weight percent Vinyi acetate, graited With 1.5 weight bercent maleic anhydride. melt index 2.0 dg/min.	P90 is Arkon P90, 10		("REB") of 90°C.
35			ol.a	de	'A 80	75	82	80	75	80	9	52	75	65	35	65	E/28VA	acet	Acid	weig	meth	etny	P90 1	Arak	("RE
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40				EX	ប	7	ຄ	7	C2	90	Ç	80	-	7	m	4									

The laminates in Table 2 were made in the same manner as in Table 1. Comparative Examples C9 and C10 are to be compared with Example 5. In Comparative Example C9, a low peel number to observed because no acid polymer is employed. In Comparative Example C10, the comonomer difference between the acid polymer and base resin polymer is greater than 10%. Comparative Example C11, which contains no tackifier resin, compares unfavorably to Example 16 which is identical in composition with the exception of the tackifier. Examples 15, 16, and 17 show that blends of otherwise identical or compatible copolymers of different melt indices can be used in place of a single copolymer.

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The laminates in Table 3 were prepared in the same manner as described for the Examples in Table 1. Comparative Examples C12 and C15 show poor results because the acrylate comonomer at 10% and vinyl ester comonomer at 9% respectively fall below the range of the present invention, i.e., 20 to 50 weight percent. Comparative Examples C13 and C14 give low peel numbers compared to Example 18 because the difference in comonomer content between acid-containing polymer and the base polymers is greater than 10%.

5		Str	g/25mm	445		-	1226	1612	1907	7.5.5 1.0.5	1651	1725	1782	1589	1930	1781	1793	2382	1532	Identified in Table I: 25 dg/min. hydrogenated C9 hydrocarbon resin, R&B 125°C. erial, R&B 140°C. hydrogenated C9 hydrocarbon resin, R&B 115°C. hydrogenated C9 hydrocarbon resin, R&B 115°C. erial, R&B 135°C. hydrogenated rosin ester from Hercules, R&B 104°C. 30% hydrogenated hydrocarbon resin from Hercules, omparable resin, is 60% hydrogenated, R&B 108°C. ated, R&B 126°C.
		א האדור ה	Д	0	0,	0.0/5	0.0	0.0	٠. د د	7.0	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	E/28VA n, R&B , R&B 1 rcules, in from
10		Adhesive mbicknoss	micrometers	31	20	31	18	20	23	20	22	25	25	25	30	25	23	30	18	id identified in Table I: d column, is identical to the first E/28VA except is 25 dg/min. 00% hydrogenated C9 hydrocarbon resin, R&B 125°C. 15% hydrogenated C9 hydrocarbon resin, R&B 115°C. material, R&B 135°C. material, R&B 135°C. 65% hydrogenated rosin ester from Hercules, R&B 104 65% hydrogenated hydrocarbon resin from Hercule a comparable resin, is 60% hydrogenated, R&B 108°C. sgenated, R&B 126°C.
15	,			15	15	15	15	15	15	15	15	15	15	15	15	15	0	15	15	al to tl hydrocar ydrocarl in ester hydroc
20			Type \$	P115	P115	P115	P90	P125	P140	F-105	R3102	R6108	R1126	M115	M135	P115	(none)	P115	F-105	in Table identical. ted C9 hydr 115°C. ted C9 hydr 115°C. ted rosin tesin, is resin, is
25	EIIa		e e		<u> </u>	.5MAn 5		20	20	20	20	20	20	20	20	20	20	20.	20	identified in Table column, is identical is 25 dg/min. hydrogenated C9 hydrerial, R&B 140°C. hydrogenated C9 hydrerial, R&B 135°C. hydrogenated rosin is comparable resin, is enated, R&B 126°C.
30	TABLE		Acid Copol	(none)	E/10iBA/10MAA	E/28VA-q-1.5MAn		•	E	t	E	2	•	=			E 000	070	20	itional materials not identified in Table I: 28vA', in the second column, is identical to the first E/28vA except that its melt index is 25 dg/min. 25 is Arkon" P125 100% hydrogenated C9 hydrocarbon resin, R&B 125·C. 40 is a comparable material, R&B 140·C. 15 is Arkon" M115 85% hydrogenated C9 hydrocarbon resin, R&B 115·C. 15 is a comparable material, R&B 135·C. 105 is Floral" 105 65% hydrogenated rosin ester from Hercules, R&B 1 102 is Regalrez" 3102, 30% hydrogenated hydrocarbon resin from Hercu R&B 102·C. R6108, a comparable resin, is 60% hydrogenated, R&B 108· R&B 102·C. R6108 a comparable resin, is 60% hydrogenated, R&B 108·
35		High m.1.	Copolymer	(none)	() II	E	E	t	=	F	2	2	E	=	=	T / AVOCA			: E	ditional materials not 1/28vA", in the second conthat its melt index is 125 is Arkon" P125 100% 140 is a comparable materials is a comparable materials is Regalrez" 3102, R&B 102°C. R6108, a control of the propersion of the prop
40			copol.	Yours of			#.	±	± 000	22.0	1		י ער פי א	G9 :		0 u	000	09	* * * *	a. Additional E/28VA', that it P125 is P P140 is 8 M115 is 8 M135 is 8 F-105 is R3102 is R&B 107 R&B 107

5	ive % MAn Peel mess in Str. meters blend g/25mm 36 0 227 33 0.4 894 41 0.3 477 38 0.3 1285 38 0.3 1407 38 0.3 1249 43 0.3 1249 51 0.03 1249 51 0.03 831 2% maleic anhydride index 1.2, grafted index 1.2, grafted in maleate terpolymer mer, melt index 5 ilymer, melt iride	
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30	High m.i. # Type # Man Peel Copolymer Acid Copol. # Type # Acid Copol. # Expons # Copolymer, melt index 1.2, grafted # Acid Copol. # Type # Acid Copolymer, melt index 2 # Expons # Copolymer, melt index 3.6, # Type # Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 43 # Type # Acid Acid Copolymer, melt index 64	
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45	E Copol. Type (none) E/28VA E/28VA E/28VA E/28VA (none) E/30NBA/10 E/30NBA/10 E/30NBA/10 E/20NBA/10	
50	EX. C12 C13 C13 C15 118 120 22 23 23	

55 methacrylic acid copolymer containing 15% methacrylic acid moieties, melt index 10 dg/min, 22% neutralized with zinc ions), 23 micrometers thick, and a layer of adhesive onto corona-treated PET film, 12 micrometers thick. The adhesive blends were melted in a 63 mm extruder with barrel temperatures from 154°C at the rear to 232°C at the front. The ionomer resin was melted in a 114 mm extruder with barrel

In Table 4, the multiple layer structures were made by coextrusion coating a layer of ionomer (ethylene

temperatures from 177 °C in the rear to 232 °C at the front. Both melt streams were fed into a 107 cm (42 inch) die set at 232 °C. The air gap was set at 16.5 centimeters. Line speed was 122 meters per minute.

Comparative Examples C17, C18, C22, C24 and C25 give poor results because the difference in vinyl ester comonomer content between the acid-containing polymer and the base polymer is greater than 10%. Comparative Example C19 compares unfavorably with Example 28 because it contains no tackifier. In Comparative Example C20, the acrylate comonomer content of the ethylene polymer is below 20 weight percent. Comparative Examples C21, C23 and C26 contain no acid polymers.

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10		Peel Str.,g/25mm greend 1 wk 4 w	423	347	276	133	428	414	450	512	563	470	510	731	557	583	509	580	290	583			•	the test sample was week and 4 weeks	
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20	Adhesive	Thickness pm	18	10.2	ω	σ	23	28	20	20	23	18	15	15	10	18	10	&	10	w	methacrylic acid copolymer, melt index 10	index 6.	110dex 25.	as soon as the test sample w conducted 1 week and 4 weeks	
		er	_				ស	ស	ນ	ß	Ŋ	ທ	15	15	15	15	15	15	15	15	nolym	melt.	mer		_
25		Tackifier Type	(none)	=	E	2	F-105	P115	R6108	E	P125	R3102	P90	F-105	P115	R6108	P90	R3102	P125	2	cid col	E/28VA, melt	,/28VA,	conduced test	nufact
		*	80	20	1 10	100	0	20	1 40	20	30	0	10	20	1 40	0	20	1 20	40	40	ic a	as E	78 E	sting o be	e E
30	TABLE IV	Copol.		10MAA	1-1.5MAr	/10MAA		10MAA	1-1.5MAr				10MAA		1-1.5MAr		10MAA	1-1.5MAr			thacry]	polymer was	the polymer was E/28VA,	to peel testing conducted wk" refer to peel testing	er sampl
35	-	Acid Cor Type	E/15MAA	E/101BA/10MAA	E/28VA-g-1.5MAn	E/10iBA/	(none)	E/10iBA/10MAA	E/28VA-g-1.5MAn	•	E	(none)	E/101BA/10MAA		E/28VA-q-1.5MAn	(none)	E/101BA/10MAA	E/28VA-g-1.5MAn	=	=		the	case the po	refers to I	respectively after sample manufacture.
40	High m.i.	_	46	0	0	o	0	0	0	15	15	15	15	15	15	0	0	0	0	0	Ethylene/15 wt. \$	each		"Green" re	respecti
45		E Coppol.b	46	80	06	•	15 6	75	53	09	40	80	09	20	30	82	65	55	45	45	à	þ.	ថ	ઌ૽	
50			C17	C18	C19	C20	C21	C22	24	25	92	C23	C24	C25	27	C26	C27	82	59	30					

In Table 5, the multiple layer structures of Examples 31 and 32 were made by coextrusion coating a layer of ionomer (as described for the Examples of Table 4), adhesive, a layer of ethylene vinyl alcohol copolymer containing 44 mol % ethylene, melt index 16, and adhesive onto PET film as described for the Examples of Table 4. The adhesive blends were melted in a 63 mm extruder with barrel temperatures from 154 °C at the rear to 232 °C at the front. The ionomer resin was melted in a 114 mm extruder with barrel

temperatures from 177 °C in the rear to 232 °C at the front. The ethylene vinyl alcohol resin was melted in a 62.5 mm extruder with barrel temperatures from 177 °C in the rear to 232 °C at the front. The melt streams were fed into a feedblock in which the adhesive stream was split into two. The 1067 mm die was at 232 °C. The air gap was set at 16.5 centimeters. Line speed was 122 meters per minute. Good adhesive strength is achieved in the structure.

The multiple layer structures of Examples 33-39 were made with several different substrates. The adhesive blends were melted in a 87.5 mm extruder with barrel temperatures from 180 °C in the rear to 240 °C at the front. The ionomer and acid copolymer resins were melted in a 62.5 mm extruder with a similar temperature profile. The melt streams were were fed into a die with a slot width of 800 mm. The die was set at 240 °C. The air gap was at 15 centimeters. Good bond strengths are obtained with different substrates.

The multiple layer structures of Examples 40-49 were prepared by coextrusion of an adhesive layer between a copolyester layer and an ethylene vinyl alcohol copolymer containing 30 mol% ethylene moieties. The copolyester was a high melt viscosity copolyester of ethylene glycol copolymerized with about 86% terephthalic and about 14% isophthalic acid, further containing 0.4% comonomer moieties derived from trimellitic acid. The adhesive blends were melted at 160 °C in a 25 mm extruder at 4-6 rpm. The copolyester was fed through a 37.5 mm extruder at 30-45 rpm at 240 °C. The extrudate was cast onto chill rolls maintained at 60 °C. The takeup speed of the sheet was 1.5 m/min. Good bonding is achieved to copolyester and ethylene vinyl alcohol copolymer with these compositions.

5				Stucture	PET/A/EVOH1/A/IONOMER1	E	PET/A/IONOMER2	8	ŧ	•	PET/A/PET	PET/A/AC/Al foil	2	POLYESTER/A/EVOH2		=					E	T	•	. Could not separate.	u		inc neutralized, 25 micrometers	ca for a sur inca		8 micr	80-120 micrometers	185-405 micrometers
15			1/25mm	6 WK		-	383	633	483	783	で	ਰ	ರ	3230	3450	4722	4859	4812	4722	4812	1816	4903	5085	l 2. d.	er (from	corona treated	neutra	25 micrometers			C # :	acids,
20			Peel Str., g/25mm	green 1 wk	583 674	574 686	367	567	450	717	p	P	p		•	!		***	1 1 1 1 1	-			!	E/28VA-q-1.5MAnh, mi	adhesive layer (from	ers, coron	224 2	10.7		ethylene,	ethylene,	trimellitic
25		ve & MAn	ss in	blend	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	9.0	9.0	0.3	0.3	9.0	0.3	0.3	0.3	9.0	E/28VA-q-		12 micrometers,	copolymen			4	HOL	and
30	>	Adhesive	H			5.1	S	10	9	10	25	5	10	9	99	99	20		55	09	09	20	22	nt 25 c.	cated, wh	e film, 1	ylic acid	opolymer.	20 micrometers	copolymer,	copolymer,	terephthalic,
35	TABLE		2 Tackifier		R3102 15	R6108 15	R3102 15	15	" 15	" 15	R6108 15	R3102 15	" 15	P125 15	" 15	٠ د	R3102 15	# 15	H 15	R3108 15	E .	n 15	15	E/28VA, mi		lyethylene terephthalate film,	15% methacr	<pre>= ec.yzene/o./* mechacryzic aczu copozymez, vlene/9% acrylic acid copolymer, melt index</pre>	511, 20 mic	alcohol		R = copolymer of iso-,
40		1. Acid		46	20	20	20	20	20	20	20	20	20	20	40	40	20	20	40	20	20	20	40	6 b.	structu	ylene to	hylene/	10 tene/	aluminum foil,	ethylene vinyl	ethylene vinyl	copolyme
4 5			S Q	40	0	0	15	15	15	15	15	15	15	15	0	15	15	0	15	15	15	0	0	E/28VA, mi	S,	lye	IONOMER1 = et	' 5	=		Z - ethyl	POLITISTER = C
50		Ethyleme	Copol.	-		2 65		50	5 50	9			9 20		1 45				5 30		ø	8 65	9 55	a. E/		PET	ONOI		A1 B	EVOL	EVOLE	POL
				EX	m	'n	'n	ñ	ñ	ň	'n	ਲ	ų.	4	4	4	4	4	4	46	4	4	4	I								

55 Claims

An extrudable bonding resin composition consisting essentially of:
 (a) 65 to 99 percent by weight of an ethylene copolymer portion consisting essentially of

(i) 1 to 100 percent by weight of a copolymer of 50 to 80 weight percent ethylene and 20 to 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and carbon monoxide, the carbon monoxide being present in an amount of 0 to 30 weight percent; having grafted thereon side chains at least one comonomer unit in length, the comonomer units in said side chains being selected from the group consisting of ethylenically unsaturated mono-, di-, or polycarboxylic acids, ethylenically unsaturated carboxylic acid anhydrides, metal salts and half-esters of such acids and anhydrides, and mixtures thereof, wherein the amount of said grafted comonomer units comprises 0.03 to 10 percent by weight of the total bonding resin composition, and

(ii) 0 to 99 percent by weight of at least one copolymer of 50 to 80 weight percent ethylene and 20 to 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and carbon monoxide, the carbon monoxide being present in an amount of 0 to 30 weight percent,

wherein the copolymer of (i) and each copolymer of (ii) contain at least one said copolymerized comonomer in common, the amount of each such common comonomer in each copolymer of (ii) being within 10% of the amount of the corresponding comonomer in copolymer (i), and the total amount of said copolymerized comonomers other than such common comonomers in any such copolymer being less than 10%; and

(b) 1 to 35 percent by weight of a tackifying resin.

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- The composition of claim 1 wherein the ethylene copolymer portion consists essentially of 5 to 50 weight percent graft copolymer of (i) and 95 to 50 weight percent of at least one copolymer of (ii).
- 3. The composition of claim 1 or 2 wherein the copolymer of (i) and each copolymer of (ii) contain the same copolymerized comonomers and each comonomer in each copolymer of (ii) is within 5% of the amount of the corresponding comonomer in copolymer (i).
 - 4. The composition of claim 1, 2 or 3 wherein the amount of ethylene copolymer portion is 80 to 95 weight percent and amount of the tackifying resin is 5 to 20 weight percent.
- The composition of any one of claims 1 to 4 wherein the amount of each copolymerized comonomer in each of the copolymers of (i) and copolymer of (ii) is 25 to 30 percent by weight.
 - 6. The composition of any one of claims 1 to 5 wherein the copolymer of (i) exclusive of the grafted comonomer units and each copolymer of (ii) are selected from the group consisting of ethylene vinyl acetate copolymer, ethylene methyl acrylate copolymer, ethylene n-butyl acrylate copolymer, and ethylene n-butyl acrylate carbon monoxide copolymer.
 - 7. The composition of any one of claims 1 to 6 wherein the at least one copolymer of (ii) comprises two copolymers which differ in melt index.
 - 8. The composition of any one of claims 1 to 7 wherein the grafted comonomer units are grafted onto polymer (i) in an amount of 0.05 to 5 weight percent of polymer (i).
- The composition of claim 8 wherein the grafted comonomer units are ethylenically unsaturated mono,
 di, or polycarboxylic acids, anhydrides, or salts.
 - 10. The composition of claim 9 wherein the anhydride is maleic anhydride.
- 11. The composition of any one of claims 1 to 10 wherein the tackifying resin has a ring and ball softening point of 0 to 150 °C and is at least one of the resins selected from the group consisting of:
 - (i) alicyclic or aliphatic hydrocarbon resins;
 - (ii) aromatic hydrocarbon resins;
 - (iii) rosin and rosin derivatives; and

(iv) terpene resins.

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- 12. A multiple layer structure comprising at least one structural layer, at least one barrier layer, and at least one bonding layer of an extrudable bonding resin composition consisting essentially of
 - (a) 65 to 99 percent by weight of an ethylene copolymer portion consisting essentially of
 - (i) 1 to 100 percent by weight of a copolymer of 50 to 80 weight percent ethylene and 20 to 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and carbon monoxide, the carbon monoxide being present in an amount of 0 to 30 weight percent; and grafted comonomer units of pendant ethylenically unsaturated mono-, di-, or polycarboxylic acids, ethylenically unsaturated carboxylic acid anhydrides, or metal salts or half-esters of such acids or anhydrides, wherein the amount of said grafted comonomer units comprises 0.03 to 10 percent by weight of the total bonding resin composition, and
 - (ii) 0 to 99 percent by weight of at least one copolymer of 50 to 80 weight percent ethylene and 20 to 50 weight percent of at least one comonomer copolymerized therewith selected from the group consisting of unsaturated mono- or dicarboxylic acids of 2-20 carbon atoms, esters or salts of said unsaturated mono- or dicarboxylic acids, vinyl esters of saturated carboxylic acids where the acid group has 1-18 carbon atoms, and carbon monoxide, the carbon monoxide being present in an amount of 0 to 30 weight percent,

wherein the copolymer of (i) and each copolymer of (ii) contain at least one said copolymerized comonomer in common, the amount of each such common comonomer in each copolymer of (ii) being within 10% of the amount of the corresponding comonomer in copolymer (i), and the total amount of said copolymerized comonomers other than such common comonomers in any such copolymer being less than 10%; and

- (b) 1 to 35 percent by weight of a tackifying resin.
- 13. The multiple layer structure of claim 12 wherein the structural layer is polyester and the barrier layer is ethylene vinyl alcohol copolymer or aluminum.
 - 14. The multiple layer structure of claim 12 comprising at least two structural layers, and wherein one structural layer is prepared from ionomer resin or acid copolymer and one structural layer is prepared from polyester.

Patentansprüche

- 1. Extrudierbare Harzbindemasse, bestehend im wesentlichen aus:
 - (a) 65 bis 99 Gew.-% eines Ethylencopolymer-Anteils, bestehend im wesentlichen aus
 - (i) 1 bis 100 Gew.-% eines Copolymeren von 50 bis 80 Gew.-% Ethylen und 20 bis 50 Gew.-% von wenigstens einem Comonomeren, das damit copolymerisiert ist, ausgewählt aus der Gruppe, bestehend aus ungesättigten Mono- oder Dicarbonsäuren mit 2 bis 20 Kohlenstoffatomen, Estern oder Salzen der genannten ungesättigten Mono- oder Dicarbonsäuren, Vinylestern von gesättigten Carbonsäuren, worin die Säuregruppe 1 bis 18 Kohlenstoffatome besitzt, und Kohlenmonoxid, wobei das Kohlenmonoxid in einer Menge von 0 bis 30 Gew.-% vorhanden ist, das darauf aufgepfropfte Seitenketten, die wenigstens eine Comonomer-Einheit ing eind, aufweist, wobei die Comonomer-Einheiten in den genannten Seitenketten ausgewählt sind aus der Gruppe, bestehend aus ethylenisch ungesättigten Mono-, Di- oder Polycarbonsäuren, ethylenisch ungesättigten Carbonsäureanhydriden, Metallsalzen und Halbestern von solchen Säuren und Anhydriden und aus Gemischen davon, worin die Menge der genannten gepfropften Comonomer-Einheiten 0,03 bis 10 Gew.-% der gesamten Harzbindemasse umfaßt, und
 - (ii) 0 bis 99 Gew.-% von wenigstens einem Copolymer und 50 bis 80 Gew.-% Ethylen und 20 bis 50 Gew.-% von wenigstens einem damit copolymerisierten Comonomer, ausgewählt aus der Gruppe, bestehend aus ungesättigten Mono- oder Dicarbonsäuren mit 2 bis 20 Kohlenstoffatomen, Estern oder Salzen der genannten ungesättigten Mono- oder Dicarbonsäuren, Vinylestern von gesättigten Carbonsäuren, worin die Säuregruppe 1 bis 18 Kohlenstoffatome besitzt, und Kohlenmonoxid, wobei das Kohlenmonoxid in einer Menge von 0 bis 30 Gew.-% vorhanden ist,

worin das Copolymer von (i) und jedes Copolymer von (ii) wenigstens ein genanntes copolymerisier-

tes Comonomer gemeinsam enthalten, wobei die Menge eines jeden solchen gemeinsamen Comonomeren in jedem Copolymer von (ii) im Bereich von 10 % der Menge des entsprechenden Comonomeren in Copolymer (i) liegt und die Gesamtmenge der genannten copolymerisierten Comonomeren, außer solchen gemeinsamen Comonomeren, in jedem solchen Copolymer weniger als 10 % beträgt, und

(b) 1 bis 35 Gew.-% eines klebrigmachenden Harzes.

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- 2. Masse nach Anspruch 1, worin der Ethylencopolymer-Anteil im wesentlichen aus 5 bis 50 Gew.-% Pfropfcopolymer von (i) und 95 bis 50 Gew.-% von wenigstens einem Copolymer von (ii) besteht,
- Masse nach Anspruch 1 oder 2, worin das Copolymer von (i) und jedes Copolymer von (ii) dieselben copolymerisierten Comonomere enthalten und jedes Comonomer in jedem Copolymer von (ii) im Bereich von 5 % der Menge des entsprechenden Comonomeren in Copolymer (i) liegt.
- Masse nach Anspruch 1, 2 oder 3, worin die Menge des Ethylencopolymer-Anteils 80 bis 95 Gew.-% beträgt und die Menge des klebrigmachenden Harzes 5 bis 20 Gew.-% beträgt.
 - 5. Masse nach einem der Ansprüche 1 bis 4, worin die Menge eines jeden copolymerisierten Comonomeren in jedem der Copolymeren von (i) und des Copolymeren von (ii) 25 bis 30 Gew.-% beträgt.
- 6. Masse nach einem der Ansprüche 1 bis 5, worin das Copolymer von (i), ausschließlich der gepfropften Comonomer-Einheiten, und jedes Copolymer von (ii) ausgewählt sind aus der Gruppe, bestehend aus einem Ethylen-Vinylacetat-Copolymer, Ethylen-Methylacrylat-Copolymer, Ethylen-n-Butylacrylat-Copolymer und einem Ethylen-n-Butylacrylat-Kohlenmonoxid-Copolymer.
 - 7. Masse nach einem der Ansprüche 1 bis 6, worin wenigstens ein Copolymer von (ii) zwei Copolymere umfaßt, die sich im Schmelzindex unterscheiden.
- 8. Masse nach einem der Ansprüche 1 bis 7, worin die gepfropften Comonomer-Einheiten auf Polymer (i) in einer Menge von 0,05 bis 5 Gew.-% von Polymer (i) gepfropft sind.
 - Masse nach Anspruch 8, worin die gepfropften Comonomer-Einheiten ethylenisch ungesättigte Mono-, Di- oder Polycarbonsäuren, Anhydride oder Salze sind.
- 10. Masse nach Anspruch 9, worin das Anhydrid Maleinsäureanhydrid ist.
 - 11. Masse nach einem der Ansprüche 1 bis 10, worin das klebrigmachende Harz einen Ring- und Ball-Erweichungspunkt von 0 bis 150 °C besitzt und wenigstens eines der Harze ist, ausgewählt aus der Gruppe, bestehend aus:
 - (i) alicyclischen oder aliphatischen Kohlenwasserstoffharzen;
 - (ii) aromatischen Kohlenwasserstoffharzen;
 - (iii) Kolophonium und Kolophonium-Derivaten; und
 - (iv) Terpenharzen.
- 45 12. Mehrschichtige Struktur, umfassend wenigstens eine Strukturschicht, wenigstens eine Barriereschicht und wenigstens eine Bindeschicht aus einer extrudierbaren Bindeharzmasse, bestehend im wesentlichen aus
 - (a) 65 bis 99 Gew.-% eines Ethylencopolymer-Anteils, bestehend im wesentlichen aus
 (i) 1 bis 100 Gew.-% eines Copolymeren von 50 bis 80 Gew.-% Ethylen und 20 bis 50 Gew.-% von wenigstens einem Comonomeren, das damit copolymerisiert ist, ausgewählt aus der Gruppe, bestehend aus ungesättigten Mono- oder Dicarbonsäuren mit 2 bis 20 Kohlenstoffatomen, Estern oder Salzen der genannten ungesättigten Mono- oder Dicarbonsäuren, Vinylestern von gesättigten Carbonsäuren, worin die Säuregruppe 1 bis 18 Kohlenstoffatome aufweist, und Kohlenmonoxid, wobei das Kohlenmonoxid in einer Menge von 0 bis 30 Gew.-% vorhanden ist, und gepfropften Comonomer-Einheiten aus angehängten ethylenisch ungesättigten Mono-, Di- oder Polycarbon-Säuren, ethylenisch ungesättigten Carbonsäureanhydriden oder Metallsalzen oder Halbestern von solchen Säuren oder Anhydriden, worin die Menge der genannten gepfropften Comonomer-Einheiten 0,03 bis 10 Gew.-% der gesamten Harzbindemasse umfaßt, und

(ii) 0 bis 99 Gew.-% von wenigstens einem Copolymer aus 50 bis 80 Gew.-% Ethylen und 20 bis 50 Gew.-% von wenigstens einem damit copolymerisierten Comonomer, ausgewählt aus der Gruppe, bestehend aus ungesättigten Mono- oder Dicarbonsäuren mit 2 bis 20 Kohlenstoffatomen, Estern oder Salzen der genannten ungesättigten Mono- oder Dicarbonsäuren, Vinylestern von gesättigten Carbonsäuren, worin die Säuregruppe 1 bis 18 Kohlenstoffatome aufweist, und Kohlenmonoxid, wobei das Kohlenmonoxid in einer Menge von 0 bis 30 Gew.-% vorhanden ist,

worin das Copolymer von (i) und jedes Copolymer von (ii) wenigstens eines der genannten copolymerisierten Comonomeren gemeinsam enthalten, wobei die Menge eines jeden solchen gemeinsamen Comonomeren in jedem Copolymer von (ii) im Bereich von 10 % der Menge des entsprechenden Comonomeren in Copolymer (i) liegt und die Gesamtmenge der genannten copolymerisierten Comonomeren, außer solchen gemeinsamen Comonomeren, in jedem derartigen Copolymer weniger als 10 % beträgt, und

- (b) 1 bis 35 Gew.-% eines klebrigmachenden Harzes.
- 13. Mehrschichtenstruktur nach Anspruch 12, worin die Strukturschicht ein Polyester und die Barriereschicht ein Ethylen-Vinylalkohol-Copolymer oder Aluminium ist.
 - 14. Mehrschichtenstruktur nach Anspruch 12, umfassend wenigstens zwei Strukturschichten, worin eine Strukturschicht aus einem Ionomerharz oder einem sauren Copolymer und eine Strukturschicht aus Polyester hergestellt ist.

Revendications

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- 1. Une composition de résine liante extrudable, constituée essentiellement de :
 - (a) 65 à 99 pour cent en poids d'un composant copolymère d'éthylène, constitué essentiellement de (i) 1 à 100 pour cent en poids d'un copolymère de 50 à 80 pour cent en poids d'éthylène et 20 à 50 pour cent en poids d'au moins un comonomère copolymérisé avec lui qui est choisi dans le groupe formé par les acides mono- ou dicarboxyliques insaturés de 2 à 20 atomes de carbone, les esters ou sels desdits acides mono- ou dicarboxyliques insaturés, les esters de vinyle d'acides carboxyliques saturés où le groupement acide compte 1 à 18 atomes de carbone, et l'oxyde de carbone, l'oxyde de carbone étant présent en une proportion de 0 à 30 pour cent en poids; sur lequel sont greffées des chaînes latérales ayant une longueur d'au moins un motif de comonomère, les motifs de comonomère desdites chaînes latérales étant choisis dans le groupe formé par les acides mono-, di- ou polycarboxyliques éthyléniquement insaturés, les anhydrides d'acides carboxyliques éthyléniquement insaturés, les sels métalliques et les hémi-esters de ces acides et anhydrides, et leurs mélanges, la quantité desdits motifs de comonomère greffé étant de 0,03 à 10 pour cent en poids de la composition de résine liante totale, et
 - (ii) 0 à 99 pour cent en poids d'au moins un copolymère de 50 à 80 pour cent en poids d'éthylène et 20 à 50 pour cent en poids d'au moins un comonomère copolymérisé avec lui qui est choisi dans le groupe formé par les acides mono- ou dicarboxyliques insaturés de 2 à 20 atomes de carbone, les esters ou sels desdits acides mono- ou dicarboxyliques insaturés, les esters de vinyle d'acides carboxyliques saturés où le groupement acide compte 1 à 18 atomes de carbone, et l'oxyde de carbone, l'oxyde de carbone étant présent en une proportion de 0 à 30 pour cent en poids,

dans lequel le copolymère de (i) et chaque copolymère de (ii) contiennent en commun au moins un comonomère copolymérisé susdit, la proportion de chaque tel comonomère commun dans chaque copolymère de (ii) ne différant pas de plus de 10 % de la proportion du comonomère correspondant dans le copolymère (i), et la proportion totale desdits comonomères copolymérisés autres que de tels comonomères communs dans chacun de ces copolymères étant inférieure à 10 % : et

- (b) 1 à 35 pour cent en poids d'une résine d'adhésivité.
- 2. La composition de la revendication 1, dans laquelle le composant copolymère d'éthylène est constitué essentiellement de 5 à 50 pour cent en poids de copolymère greffé de (i) et 95 à 50 pour cent en poids d'au moins un copolymère de (ii).
 - 3. La composition de la revendication 1 ou 2, dans laquelle le copolymère de (i) et chaque copolymère de (ii) contiennent les mêmes comonomères copolymérisés et la proportion de chaque comonomère dans

chaque copolymère de (ii) ne diffère pas de plus de 5 % de la proportion du comonomère correspondant dans le copolymère (i).

- 4. La composition de la revendication 1, 2 ou 3, dans laquelle la quantité du composant copolymère d'éthylène est de 80 à 95 pour cent en poids et la quantité de la résine d'adhésivité est de 5 à 20 pour cent en poids.
 - 5. La composition de l'une quelconque des revendications 1 à 4, dans laquelle la proportion de chaque comonomère copolymérisé dans chacun des copolymères de (i) et du copolymère de (ii) est de 25 à 30 pour cent en poids.
 - 6. La composition de l'une quelconque des revendications 1 à 5, dans laquelle le copolymère de (i) à l'exclusion des motifs de comonomère greffé et chaque copolymère de (ii) sont choisis dans le groupe formé par un copolymère éthylène-acétate de vinyle, un copolymère éthylène-acrylate de méthyle, un copolymère éthylène-acrylate de n-butyle et un copolymère éthylène-acrylate de n-butyle-oxyde de carbone.
 - 7. Le composition de l'une quelconque des revendications 1 à 6, dans laquelle ledit au moins un copolymère de (ii) comprend deux copolymères qui diffèrent par l'indice de fluidité à l'état fondu.
 - 8. La composition de l'une quelconque des revendications 1 à 7, dans laquelle les motifs de comonomère greffé sont greffés sur le polymère (i) en une proportion de 0,05 à 5 pour cent en poids du polymère (i).
- 9. La composition de la revendication 8, dans laquelle les motifs de comonomère greffé sont des acides, anhydrides ou sels mono-, di- ou polycarboxyliques éthyléniquement insaturés.
 - 10. La composition de la revendication 9, dans laquelle l'anhydride est l'anhydride maléique.
- 11. La composition de l'une quelconque des revendications 1 à 10, dans laquelle la résine d'adhésivité a un point de ramollissement de 0 à 150°C par la méthode bille et anneau et est au moins l'une des résines choisies dans le groupe formé par :
 - (i) les résines hydrocarbonées alicycliques ou aliphatiques ;
 - (ii) les résines hydrocarbonées aromatiques ;
 - (iii) la colophane et les dérivés de la colophane ; et
 - (iv) les résines terpéniques.

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- 12. Une structure multicouche comprenant au moins une couche structurale, au moins une couche de barrière et au moins une couche de liaison formée d'une composition de résine liante extrudable constituée essentiellement de
 - (a) 65 à 99 pour cent en poids d'un composant copolymère d'éthylène, constitué essentiellement de (i) 1 à 100 pour cent en poids d'un copolymère de 50 à 80 pour cent en poids d'éthylène et 20 à 50 pour cent en poids d'au moins un comonomère copolymérisé avec lui qui est choisi dans le groupe formé par les acides mono- ou dicarboxyliques insaturés de 2 à 20 atomes de carbone, les esters ou sels desdits acides mono- ou dicarboxyliques insaturés, les esters de vinyle d'acides carboxyliques saturés où le groupement acide compte 1 à 18 atomes de carbone, et l'oxyde de carbone, l'oxyde de carbone étant présent en une proportion de 0 à 30 pour cent en poids ; et de motifs de comonomère greffé formés par des acides mono-, di- ou polycarboxyliques éthyléniquement insaturés, ou des sels métalliques ou hémi-esters de ces acides et anhydrides, en situation latérale, la quantité desdits motifs de comonomère greffé représentant 0,03 à 10 pour cent en poids de la composition de résine liante totale, et
 - (ii) 0 à 99 pour cent en poids d'au moins un copolymère de 50 à 80 pour cent en poids d'éthylène et 20 à 50 pour cent en poids d'au moins un comonomère copolymérisé avec lui qui est choisi dans le groupe formé par les acides mono- ou dicarboxyliques insaturés de 2 à 20 atomes de carbone, les esters ou sels desdits acides mono- ou dicarboxyliques insaturés, les esters de vinyle d'acides carboxyliques saturés où le groupement acide compte 1 à 18 atomes de carbone, et l'oxyde de carbone, l'oxyde de carbone étant présent en une proportion de 0 à 30

pour cent en poids,

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dans lequel le copolymère de (i) et chaque copolymère de (ii) contiennent en commun au moins un comonomère copolymérisé susdit, la proportion de chaque tel comonomère commun dans chaque copolymère de (ii) ne différant pas de plus de 10 % de la proportion du comonomère correspondant dans le copolymère (i), et la proportion totale desdits comonomères copolymérisés autres que de tels comonomères communs dans chacun de ces copolymères étant inférieure à 10

- (b) 1 à 35 pour cent en poids d'une résine d'adhésivité.
- 13. La structure multicouche de la revendication 12, dans laquelle la couche structurale est formée d'un polyester et la couche de barrière est formée d'un copolymère éthylène-alcool vinylique ou d'alumi-
- 14. La structure multicouche de la revendication 12, comprenant au moins deux couches structurales, et dans laquelle une couche structurale est préparée à partir de résine ionomère ou de copolymère acide, 15 et une couche structurale est préparée à partir de polyester.